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The intracellular immune receptor Rx1 regulates the DNA-binding activity of a Golden2-like transcription factor*

Philip D. Townsend^{1,2,7}, Christopher H. Dixon^{1,2,7}, Erik J. Sloatweg⁴, Octavina C.A. Sukarta⁴, Ally W.H. Yang⁵, Timothy R. Hughes⁵, Gary J. Sharples^{1,2}, Lars-Olof Pålsson³, Frank L.W. Takken⁶, Aska Goverse⁴, and Martin J. Cann^{1,2}

¹Department of Biosciences, ²Biophysical Sciences Institute, ³Department of Chemistry, Durham University, South Road, Durham, DH1 3LE, United Kingdom.

⁴Laboratory of Nematology, Department of Plant Sciences, Wageningen University, 6708 PB, Wageningen, The Netherlands.

⁵Donnelly Centre, University of Toronto, 160 College Street, Toronto, ON, M5S 3E1, Canada

⁶Molecular Plant Pathology, Swammerdam Institute for Life Sciences, University of Amsterdam, Science Park 904, 1098 XH, Amsterdam, The Netherlands.

⁷ These authors contributed equally to this work

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To whom correspondence should be addressed: Martin J. Cann, Department of Biosciences, Durham University, South Road, Durham, DH1 3LE, United Kingdom. Phone: +44 (191) 3343985. Fax: +44 (191) 3341201. E-mail: m.j.cann@durham.ac.uk.

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ABSTRACT.

Plant NLR proteins enable the immune system to recognise and respond to pathogen attack. An early consequence of immune activation is transcriptional reprogramming and some NLRs have been shown to act in the nucleus and interact with transcription factors. The Rx1 NLR protein of potato is further able to bind and distort double-stranded DNA. However, Rx1 host targets that support a role for Rx1 in transcriptional reprogramming at DNA are unknown. Here we report a functional interaction between Rx1 and NbGlk1, a Golden2-like transcription factor. Rx1 binds to NbGlk1 *in vitro* and *in planta*. NbGlk1 binds to known Golden2-like consensus DNA sequences. Rx1 reduces the binding affinity of NbGlk1 for DNA *in vitro*. NbGlk1 activates cellular responses to potato virus X, whereas Rx1 associates with NbGlk1 and prevents its assembly on DNA *in planta* unless activated by PVX. This study provides new mechanistic insight into how an NLR can co-ordinate an immune signalling response at DNA following pathogen perception.

Plants possess an innate immune system that enables cell-autonomous defence responses upon pathogen perception (1,2). Plant NLR⁸ immune receptors detect strain-specific pathogen effectors to mediate the immune responses to an invading pathogen (2-4). NLR proteins belong to the STAND P-loop ATPases of the AAA superfamily, and have a multidomain structure that allows them to function as a sensor, switch and response factor (5,6).

The NLR N-terminus typically consists of either a CC or TIR domain (7). The NLR NB domain, also referred to as the NB-ARC domain, is proposed to function as a molecular switch in NLR activation (6,8-10). The C-terminal LRR domain is required for pathogen recognition specificity and for maintaining the NLR protein in an auto-inhibited state. The NB-ARC domain of tomato I-2 and Mi-1, flax M and L6, and barley MLA27 is ADP-bound in this auto-inhibited state (7,11,12). Pathogen recognition via the LRR domain is proposed to permit the exchange of ADP for ATP, allowing the NB-ARC domain to adopt its activated state. ATP hydrolysis to ADP is proposed to re-establish the

inactivated state. For example, mutants of the tomato I-2 NLR with reduced levels of *in vitro* ATP hydrolysis are auto-activated *in vivo* (11). Nucleotide hydrolysis in NLR inactivation may extend further than ADP. For example, the NB subdomain of rice Os2g_25900 and the NB-ARC domains of maize PSiP and *Arabidopsis* Rpm1 possess a nucleotide phosphatase activity in which all phosphates are removed from the nucleotide triphosphate to leave the nucleoside base (13).

A crucial question concerns the nature of the downstream signalling component(s) for plant NLR proteins and how these are activated or inactivated by NLR proteins upon pathogen perception. Several NLR proteins, including N, MLA10, and Rx1 have a dynamic nuclear-cytoplasmic distribution, whereas RRS1-R is restricted to the nucleus, dependent upon the presence of the PopP2 immune elicitor (14-18). Several NLRs, including, barley MLA1 and MLA10, *Arabidopsis* RPS4 and SNC1, and the tobacco N protein show a nuclear localization (17,19-21). Redirection of MLA10, N, RPS4, and SNC1 from the nucleus to the cytoplasm reduces immune activation suggesting a signalling component resident in the nucleus (14,17,20,22). One of the most important and earliest consequences of immune activation is transcriptional reprogramming (23-25). The association of MLA10 with Myb and WRKY transcription factors suggests that plant NLRs themselves might regulate transcription in the immune response (26,27).

Biochemical data suggests that at least a subset of plant NLRs are directly active at DNA. Rx1 of potato, I-2 of tomato, and the orphan NLR PSiP of maize, interact directly with DNA *in vitro* (28,29). The *Rx1* gene, introgressed in potato from the wild species *Solanum tuberosum* subsp. *andigena*, confers resistance to PVX upon recognition of its coat protein (30,31). The Rx1 protein binds to genomic DNA *in situ* dependent upon immune activation (29). In addition, Rx1 induces ATP-dependent bending and melting of DNA *in vitro*. Analysis of Rx1-binding to a variety of DNA structures demonstrated a preference for topologies resembling transcription bubbles. Rx1 therefore binds, bends and distorts DNA in a manner reminiscent of the formation of the transcription initiation complex (32-35).

In vitro analysis demonstrated no sequence specificity in the Rx1 interaction with dsDNA. There is a question, therefore, over how the non-specific interaction of Rx1 with DNA can be reconciled with a specific role in immune activation. The Rx1 CC domain is responsible for the nuclear accumulation of Rx1. Furthermore, the Rx1 CC

domain diffusion rate in the nucleus is low, pointing to complex formation with nuclear components (18). We therefore set out to identify nuclear interactors of the Rx1 CC domain and investigate their role with Rx1 in immunity at DNA. Here we demonstrate that Rx1 interacts directly with a Golden2-like TF (*NbGlk1*). This Golden2-like TF mediates immune responses to PVX and has its activity at DNA regulated by Rx1. The findings therefore provide new insight into the action of NLR proteins at genomic DNA in controlling immune responses.

RESULTS

The Rx1 CC domain interacts with a DNA associated protein - We set out to identify Rx1 CC domain interactors and investigate their role in immunity at DNA. Amino acids 1-144 of Rx1, encompassing the CC domain, were used as bait in a Yeast Two-Hybrid screen using a random-primed *N. benthamiana* mixed tissue cDNA library. We identified a total of six clones with sequence similarity to a GLK TF (here called *NbGlk1*) (Figure 1). The six clones corresponded to three individual cDNAs isolated twice each. Individual clones were presumably obtained multiple times due to amplification of the cDNA library. The full-length *NbGlk1* cDNA (Figure S1A) encodes a 402-amino acid protein of predicted molecular weight 44531 Da and carries a single Myb-type helix-turn-helix DNA-binding domain. GLK TFs are classified into the GARP TF family (36). *NbGlk1* possessed a C-terminal GCT box specific to GLK-type TFs and a AREA₂EAA hexapeptide sequence within the DNA-binding domain typical of GARP family TFs (37,38) (Figure S1B).

Rx1 interacts directly with NbGlk1 in vitro and in vivo - The *NbGlk1* Yeast Two-Hybrid clones all encompassed the Myb-type DNA-binding domain. We therefore assessed Rx1-binding to *NbGlk1* before analysing their interactions with DNA. Unfortunately, we were unable to express full-length *NbGlk1* as a recombinant protein. We therefore expressed amino acids 83-402 of *NbGlk1* (*NbGlk1*(83-402)) and examined its interaction with the Rx1-CC domain (Rx1(1-144)) by size exclusion chromatography. Rx1(1-144) ran predominantly in the void volume (Figure 2A, upper SDS-PAGE panel, elution volume of 35-37 mLs). We hypothesise that this peak consists of a higher order oligomeric Rx1-CC complex but its relevance to Rx1 biochemistry, if any, is not known. We noted a shift in the peak bands corresponding to Rx1(1-144) (Figure 2A, upper SDS-PAGE panel, capped green

bar) and *NbGlk1*(83-402) (Figure 2A, middle SDS-PAGE panel, capped red bar) when incubated together (Figure 2A, lower SDS-PAGE panel) showing that *NbGlk1* and Rx1-CC interact directly *in vitro*.

It is possible that the observed interaction between Rx1(1-144) and *NbGlk1*(83-402) was influenced by an unknown property of Rx1(1-144) that also causes a proportion of the protein to run in the gel filtration void volume. To enhance protein solubility we developed a new Rx1-CC protein with an *N*-terminal GST tag. We assessed binding of the Rx1-CC domain-GST fusion (Rx1(GST-1-144) to *NbGlk1*(83-402). Rx1(GST-1-144) was completely soluble with no detectable protein in the void volume (Figure 2B, UV trace). However, Rx1(GST-1-144) was susceptible to some proteolytic cleavage during purification at the extreme *C*-terminus and resulted in the protein running as a doublet by SDS-PAGE. We noted a shift in the peak bands corresponding to Rx1(GST-1-144) (Figure 2B, SDS-PAGE panel 1, capped green bar) and *NbGlk1*(83-402) (Figure 2B, SDS-PAGE panel 2, capped red bar) when incubated together (Figure 2B, SDS-PAGE panel 3). GST protein expressed alone (Figure 3, SDS-PAGE panel 4) showed no shift in peak band when expressed with *NbGlk1*(83-402) (Figure 2B, SDS-PAGE panel 5). *NbGlk1* and Rx1(GST-1-144) therefore interact directly *in vitro*.

We next assessed whether full-length Rx1 and *NbGlk1* interact *in planta*. We performed co-immunoprecipitation experiments using full-length *NbGlk1* fused to a 4xHA epitope tag with either full-length Rx1 or the CC domain fused to a 4xMyc epitope tag. *NbGlk1*-4HA, when co-expressed with full-length Rx1 or the CC domain, was immunoprecipitated using the anti-myc antibody and could be detected with the anti-HA tag antibody (Figure 3). A very faint immunoblot band for *NbGlk1*-4HA was observed after immunoprecipitation using an anti-myc antibody when co-expressed with a fusion of GFP to a 4xmyc epitope tag. This signal was significantly lower than those obtained on co-expression with Rx1 indicating that its immunoprecipitation was specific. Full-length *NbGlk1* therefore interacts with the CC domain and full-length Rx1 *in planta*.

Rx1 modulates NbGlk1 interactions with DNA in vitro - We investigated whether *NbGlk1* showed sequence-specific DNA-binding. We analysed the DNA-binding properties of the *NbGlk1* DNA-binding domain compared to the known binding properties of GLK TFs using a protein-binding microarray (PBM) consisting of ~41,000 35-mer

probes in which all possible 10-mers occur once and all nonpalindromic 8-mers are represented 32 times, allowing for an unbiased assessment of sequence preference for all possible 8-mers (39). Values for individual 8-mers were obtained as *E* scores (representing relative rank of intensities ranging from -0.5 to + 0.5) and *Z* scores (scaling approximately with binding affinity) (Figure S2). The DNA-binding sequence for the top scoring 8-mer for *NbGlk1* was 5'-AGATTTCC-3' and 5'-AGATTTCC-3' for *E* score (0.49648) and *Z* score (37.25190), respectively. Both identified DNA motifs are similar to the AGATTCT core palindrome recognised by the GLK TF encoded by At2g20570 of *Arabidopsis thaliana* (40).

We measured the K_d of *NbGlk1* for dsDNA substrates lacking the hypothesised *NbGlk1*-binding site (No site), with a concatenated AGATTTCC binding site (from the *NbGlk1* PBM, labelled AGATTT) and with a concatenated GGATATCC binding site (from the GLK1 consensus site (40) and also identified in the PBM analysis of *NbGlk1* (*Z* score=29.62117, labelled GATATC) by fluorescence anisotropy (Table 1). A recombinant protein consisting of *NbGlk1* residues 1-243 (*NbGlk1*(1-243)) encompassing the Myb-type helix-turn-helix DNA-binding domain displayed only a weak affinity ($K_d > 1 \mu\text{M}$; Table 1) for the GATATC binding site (Figure 4A). This low affinity prevented drawing conclusions on the influence of Rx1 on *NbGlk1*(1-243) DNA-binding (Figure 4A, Table 1).

Analysis of the *NbGlk1* ORF revealed a pI of 3.8 for the *N*-terminal 82 amino acids. We hypothesised that the acidic *N*-terminal 82 amino acids could interfere with analysis of the interactions of the *NbGlk1* Myb-type domain with DNA. We therefore measured *NbGlk1* DNA-binding with a protein lacking the *N*-terminal 82 amino acids. *NbGlk1*(83-402) possessed a similar K_d for both DNA containing no *NbGlk1*-binding site (Figure 4B) or a AGATTT binding site (F-test, $p=0.41$) (Figure 4C). However, *NbGlk1*(83-402) possessed a lower K_d for the GATATC binding site (F-test, $p<0.0065$) (Figure 4D). In comparison to *NbGlk1*(83-402), a C-terminally truncated recombinant protein consisting of amino acids 83-243 of *NbGlk1* (*NbGlk1*(83-243)) showed a similar K_d for DNA irrespective of whether it contained no *NbGlk1*-binding site (Figure 4E) or the AGATTT site (F-test, $p=0.507$) (Figure 4F) or GATATC sites (F-test, $p=0.0827$) (Figure 4G). There was also no significant difference in affinity between *NbGlk1*(83-402) and *NbGlk1*(83-243) for the No site ($p=0.3659$) and AGATTT ($p=0.2171$) DNA-binding motifs. However, there was a significant

difference in affinity between *NbGlk1*(83-402) and *NbGlk1*(83-243) for the GATATC ($p < 0.001$) DNA-binding motif. In conclusion, *NbGlk1* shows a higher affinity for DNA carrying a GLK family binding site than for a random sequence. Further, both the *N*- and *C*-termini of the protein contribute to DNA-binding affinity. The acidic *N*-terminal 82 amino acids significantly reduced DNA-binding. Amino acids 243-402 also enhance affinity for the GATATC motif compared to the No site DNA. Further, amino acids 243-402 enhance affinity for all DNA sequences examined. For example, *NbGlk1*(83-402) showed a higher affinity than *NbGlk1*(83-243) for No site DNA (F-test, $p < 0.001$), the AGATTT motif (F-test, $p = 0.0196$) and the GATATC motif (F-test, $p = 0.0436$). We therefore hypothesize that a region(s) within amino acids 243-402 contributes to DNA-binding.

Since our underlying hypothesis was that *NbGlk1* interacted with Rx1 at DNA, we investigated the influence of amino acids 1-489 of Rx1 (consisting of the CC and NB-ARC domains; Rx1(1-489)) on *NbGlk1* dsDNA binding affinity. Experiments were performed using conditions under which only *NbGlk1* and not Rx1 contributed to DNA-binding. These conditions were established by using a concentration of Rx1(1-489) that gave no change in anisotropy when incubated with DNA alone. *NbGlk1*(83-402) and *NbGlk1*(83-243) showed a higher K_d for DNA containing no *NbGlk1*-binding site (Figure 4A, D), the AGATTT binding site (Figure 4B, E) and the GATATC binding sites (Figure 4C, F) in the presence of Rx1(1-489) (F-test, $p < 0.0001$). Similar data was observed for the influence of Rx1(1-144) (CC domain only) on *NbGlk1*(83-402) and *NbGlk1*(83-243) DNA-binding (Table 1, Figure 5A-G) (F-test, $p < 0.0001$). We confirmed that the increased K_d was specific for *NbGlk1* by using the CAP TF of *E. coli*, which has a similar MW and pI to *NbGlk1*, as a negative control. Rx1(1-144) reduced *NbGlk1* but not CAP-binding to dsDNA (Figure 4H). ATP stimulates Rx1(1-489)-mediated dsDNA distortion (29). We therefore investigated whether ATP and/or ADP modified the influence of Rx1(1-489) on *NbGlk1* DNA-binding. We observed no effect indicating that the switch function of the NB-ARC domain does not influence *NbGlk1* DNA-binding (Figure 5I). In conclusion, the Rx1-CC and -CCNB-ARC domains reduce the affinity of *NbGlk1* for DNA. The data is consistent with the observation that the Rx1 CC binding surface overlaps with the *NbGlk1* DNA-binding domain. Rx1 may therefore restrict access of *NbGlk1* to DNA whether binding is to a consensus or non-consensus binding site.

Although both Rx1(1-489) and Rx1(1-144) reduce the binding affinity of *NbGlk1* for DNA, a potential alternative explanation for the reduction in *NbGlk1* DNA-binding affinity caused by Rx1(1-144) is that the higher ordered Rx1-CC complex observed by gel filtration analysis (Figure 2A) is sufficient to reduce the concentration of soluble *NbGlk1*. This might therefore manifest as an apparent reduction in DNA-binding affinity. We therefore assessed the influence of Rx1(GST-1-144) on *NbGlk1*(83-402) binding to DNA containing no *NbGlk1*-binding site (Figure 6A), the AGATTT site (Figure 6B) or GATATC site (Figure 6C). *NbGlk1*(83-402) showed a higher K_d for DNA containing no *NbGlk1*-binding site (Figure 6A), the AGATTT binding site (Figure 6B) and the GATATC binding sites (Figure 6C) in the presence of Rx1(GST-1-144) (F-test, $p < 0.0001$) (Table 2). The Rx1 CC domain therefore reduces the affinity of *NbGlk1*(83-402) for DNA and this observation is not an artefact due to Rx1(1-144) complex formation.

Rx1 and NbGlk1 interact at DNA in situ - We next investigated Rx1 and *NbGlk1* interactions with DNA in the plant cell. We first used confocal scanning laser microscopy to confirm the distribution of *NbGlk1* to the nucleus. The subcellular localization of *NbGlk1* *in planta* was determined by transiently expressing *NbGlk1*-GFP in the epidermal cells of *N. benthamiana*. *NbGlk1*-GFP was expressed either with or without the tombusvirus p19 silencing suppressor to enhance protein accumulation. In line with its role as a transcription factor, strong fluorescence of *NbGlk1*-GFP was observed exclusively in the nucleus (Figure 7A). This pattern does not resemble free mCherry, which has a nucleocytoplasmic distribution. The pattern of *NbGlk1*-GFP localization was not affected by p19 overexpression, minimizing the likelihood an overexpression artefact. Upon closer examination, the fluorescence of *NbGlk1*-GFP was restricted to the nucleoplasm, with a lack of signal in the nucleolus. A very small amount of GFP signal was observed in the nucleolus in the absence of p19. This was most likely background due to the longer opening of the pinhole of the microscope (pinhole of 1 for imaging with p19 and pinhole of 1.5 without p19).

Having established that *NbGlk1* is localized exclusively in the nucleus, we studied Rx1-*NbGlk1*-DNA interactions using Förster resonance energy transfer-fluorescence lifetime imaging microscopy. FRET-FLIM has been used previously to demonstrate Rx1 binding to DNA in response to

immune activation (29). GFP (negative control), histone H2B fused to GFP (GFP-H2B; positive control), full-length Rx1 with or without an N-terminal GFP tag, or full-length *NbGlk1* with or without an N-terminal GFP tag were transiently expressed in *N. benthamiana*. The constituent fluorescence lifetimes of the GFP tag were examined in leaves counterstained with the nucleic acid stain LDS 751. GFP showed two distinct lifetimes at ~0.5 and ~1.5 ns. Different extents of energy transfer from GFP to acceptor LDS 751 modulates the relative contribution of these two lifetimes. A drop in the ratio of the ~1.5 (long) to ~0.5 (short) ns GFP lifetimes indicates an interaction with DNA in the cell (29). First, we monitored the interaction of an Rx1-GFP fusion with DNA with or without *NbGlk1* (untagged) in the presence or absence of the avirulent PVX coat protein (CP106). Rx1-GFP expressed without *NbGlk1* only bound DNA in the presence of CP106 as expected (29) (Figure 7B). In contrast, Rx1-GFP co-expressed with untagged *NbGlk1* bound DNA irrespective of CP106 presence. Overexpressed *NbGlk1* is therefore able to recruit Rx1 to DNA. Second, we monitored the interaction of a *NbGlk1*-GFP tag with DNA with or without Rx1 (untagged) in the presence or absence of CP106. Surprisingly, *NbGlk1*-GFP only bound DNA in the joint presence of Rx1 and CP106 (Figure 7C). We noted variation in the ratio of lifetime yields for GFP between the two sets of experiments (Figure 7B-C). This variation might have a biological (e.g. time of year for sampling and expression) or technical (e.g. fitting) cause. Regardless, the value for the ratio of lifetimes for GFP is distinct from the positive control within and between experiments.

NbGlk1 reduces susceptibility to PVX - We hypothesized that *NbGlk1* plays a role in Rx1-mediated immunity as it interacts with DNA in response to PVX. We used virus-induced gene silencing (41,42) to investigate the *NbGlk1* requirement for Rx1-mediated immunity. VIGS with independent *NbGlk1* clones gave a bleaching phenotype consistent with a previously described role in the formation of the photosynthetic apparatus (43). The silenced leaves were too fragile for further infiltration and precluded an analysis of immune function by this method. We therefore used *NbGlk1* over-expression to investigate a role in immunity.

Rx1 mediates two forms of immunity: a) a cell death response promoted by expression of CP106 and b) symptomless virus resistance (called extreme resistance) (30). We first investigated a role for *NbGlk1* in CP106-mediated cell death. CP106-

mediated cell death is activated through the sole expression of the PVX CP106 coat protein in the absence of other viral proteins. *NbGlk1*, Rx1 and CP106 were transiently expressed in *N. benthamiana*. Rx1-mediated immune responses were assessed by scoring cell death within the infiltrated areas. Cell death associated with the hypersensitive response was observed in the presence of Rx1 and CP106 and was not influenced by *NbGlk1* (Figure 8A-C). Cell death was not as extensive as previously reported for Rx1 and CP106 co-expression (18) as agroinfiltration was performed at the same low $A_{600\text{ nm}}$ that was used for the FRET-FLIM analysis (Figure 7B-C).

Next, we investigated a role for *NbGlk1* in PVX-mediated extreme resistance. PVX-mediated extreme resistance is activated by the virus following detection of CP106. *NbGlk1*, Rx1 and a PVX amplicon in which GFP expression is driven from a coat protein promoter were transiently expressed in *N. benthamiana*. The Rx1 expression construct possessed an out-of-frame second start codon introduced upstream of the genuine Rx1 start codon reducing translation efficiency and increasing the sensitivity of the assay to immune modifiers (18). We monitored virus resistance by visualizing GFP expression within the infiltrated region of the leaf. Rx1 suppressed PVX-directed GFP expression as expected (Figure 8D). Surprisingly, overexpressed *NbGlk1* was sufficient to suppress PVX-directed GFP expression independent of Rx1. Rx1 co-expression with *NbGlk1* did not further reduce GFP levels. Over-expressed *NbGlk1* is therefore able to bypass the requirement for Rx1 in extreme resistance to PVX.

DISCUSSION

A Yeast Two-Hybrid screen identified the GLK-like TF, *NbGlk1*, as an Rx1-interacting protein (Figure 1, 2, 3). GLK-like TFs are involved in defence signalling in *Arabidopsis*, providing resistance to cucumber mosaic virus (44) and the fungal pathogens *Fusarium graminearum* (45) and *Hyaloperonospora arabidopsidis* (46). *NbGlk1* therefore links Rx1 with a TF class known to function in biotic stress (47). Rx1 binds and distorts DNA *in vitro* but without an apparent sequence specificity (29). *NbGlk1* could therefore provide the sequence selectivity for Rx1 at DNA. *NbGlk1* bound dsDNA non-specifically but showed a higher affinity for specific DNA motifs bound by the related GLK1 TF (At2g20570 of *A. thaliana*) (40). TFs generally show a lower affinity for non-specific DNA sequences (48) and this may assist TFs to scan

DNA in searches for their cognate motifs (49). *NbGlk1* therefore has the properties of a TF that can target Rx1 to a specific DNA motif.

The *NbGlk1* N-terminus reduced the affinity for DNA *in vitro* implying an autoinhibitory role of this domain in DNA-binding (Figure 4A, Table 1). In agreement, full-length *NbGlk1* over-expressed in the plant does not show an interaction with DNA by FRET-FLIM (Figure 7C). Co-incubation of Rx1 with *NbGlk1* decreases the affinity of *NbGlk1* for DNA *in vitro* (Figure 4B-G, 5A-G, Figure 6A-C, Table 1 & 2). We propose this arises from the Rx1-CC-binding surface overlapping the *NbGlk1* DNA-binding domain.

NbGlk1 is not bound to DNA *in planta* unless Rx1 is activated through PVX-derived CP106 (Figure 7C). Rx1 does not interact with DNA unless it is activated by CP106 or when co-expressed with *NbGlk1* *in planta* (Figure 7B). A caveat with the interpretation of the FRET-FLIM data is the possibility of false negative results. If the expressed GFP fusion protein has saturated all available DNA-binding sites, the accumulation of an increased pool of non-DNA-bound protein will shift the ratio of the long to short lifetimes to the GFP negative control. In the current analysis, however, the *in vitro* data is entirely consistent with the FRET-FLIM analysis. In the absence of an available alternative method not susceptible to the same issue of false negatives, however, the interpretation of the data should be viewed with some caution.

One interpretation of these data is that co-expression of Rx1 and *NbGlk1* permits complex formation at DNA. However, this complex might form at amounts escaping the detection level of the FRET-FLIM assays (Figure 9A, right hand side). Following co-expression of Rx1 and *NbGlk1*, the DNA-bound state of the complex might be stabilized by the combined intrinsic DNA binding activity of Rx1 with the low affinity of *NbGlk1* for non-consensus sequences. The joint affinity of both proteins for DNA shifts the equilibrium to a DNA-bound state detectable in our setup (Figure 9A, left hand side). The Rx1-*NbGlk1* complex could be arranged such that specifically Rx1 contacts DNA and the auto-inhibited *NbGlk1* stabilizes this complex by its weak interaction with non-consensus sequences. Immune activation via PVX permits an uncharacterized structural change in the complex releasing the negative regulation on *NbGlk1*, permitting it to identify and directly and strongly interact with its consensus sequence (Figure 9B, right hand side). Once *NbGlk1* is stably bound to its consensus sequences immune signaling is activated. This model is in agreement with the observation that

overexpression of *NbGlk1* alone overcomes the need for Rx1; apparently in this situation a sufficient large pool of active *NbGlk1* is present to identify and interact to these consensus motifs and to trigger immunity. Over-expressed *NbGlk1* was not observed to bind DNA *in planta* in our assays, but it should be noted that the fluorescent techniques used to monitor DNA-binding (FRET-FLIM) are less sensitive than those monitoring immunity (steady state fluorescence). Over-expressed *NbGlk1* complex will be in a thermodynamically-coupled cycle of four states; non-DNA-bound or DNA-bound at either a non-consensus site or a *NbGlk1* consensus site. Hence, even if the bound state at a consensus site is thermodynamically non-favored in the absence of Rx1/CP106 sufficient binding may occur on over-expression to permit an immune response.

The function of Rx1 may therefore be to enable the specific activation of *NbGlk1* in response to PVX by releasing auto-inhibition allowing *NbGlk1* to scan for, and interact, with its consensus sequences *in planta*. In conclusion, we identify *NbGlk1* as an immune activating protein acting at DNA and regulated by Rx1. These observations provide a direct and unexpected link between NLR-mediated perception of PVX and transcriptional processes at DNA. More generally, the findings suggest that nuclear-localised TFs involved in immunity are inactive until de-repressed by an activated NLR following pathogen perception.

EXPERIMENTAL PROCEDURES

Oligonucleotides – All oligonucleotide sequences used for this study are provided in Supplementary Table 1.

Plasmids – The *NbGlk1* (Niben101Scf06721g00011.1 (<https://solgenomics.net>) open-reading frame was amplified by PCR from cDNA synthesized from *Nicotiana benthamiana* whole leaf material and the DNA sequenced on both strands by Sanger DNA sequencing (Figure S1A). Several differences from the computed open-reading frame for Niben101Scf06721g00011.1 were noted. The cloned open-reading frame is shown in Figure S1B. A PCR product spanning *NbGlk1* residues 83-402 of an *E. coli* codon optimized synthetic cDNA (Genscript, Figure S1C) was cloned into the NdeI and XhoI sites of pET14b (pET14b-*NbGlk1*(83-402) and fitted with a hexahistidine tag for affinity purification of recombinant protein. The oligonucleotides used to construct pET14b-*NbGlk1*(83-402) were *NbGlk1*-1 and *NbGlk1*-2. Similarly, PCR products spanning

residues 83-243 and 1-243 of the *NbGlk1* synthetic cDNA were cloned into the *NdeI* and *BamHI* sites of pET14b (pET14b-*NbGlk1*(83-243) and pET14b-*NbGlk1*(1-243)) and fitted with hexahistidine tags for affinity purification of recombinant protein. The oligonucleotides used to construct pET14b-*NbGlk1*(83-243) were *NbGlk1*-3 and *NbGlk1*-4. The oligonucleotides used to construct pET14b-*NbGlk1*(1-243) were *NbGlk1*-5 and *NbGlk1*-6. A PCR product spanning residues 82-244 of the *NbGlk1* synthetic cDNA was cloned into the *AscI* and *SbfI* sites of pTH6838, a modified T7-driven GST expression vector (pTH6838-*NbGlk1*(82-244)) (39).

A PCR product spanning residues 1-144 of Rx1 (GenBankTM accession number AJ011801.1) was cloned into the *NcoI* and *NotI* sites of pET28a (pET28a-Rx1-CC) and fitted with a hexahistidine tag for affinity purification of recombinant protein. The oligonucleotides used to construct pET28a-Rx1-CC were Rx1-1 and Rx1-2. A PCR product spanning residues 1-489 of Rx1 was cloned into the *NdeI* and *XhoI* sites of pET22b (pET22b-Rx1-CCNBARC) to give an N-terminal hexahistidine tag for affinity purification of recombinant protein. The oligonucleotides used to construct pET22b-Rx1-CCNBARC were Rx1-3 and Rx1-4. A PCR product spanning Rx1 residues 1-144 was cloned into the *BamHI* and *XhoI* sites of pGEX-6P-1 (pGEX-6P-1-Rx1(1-144)) and fitted with a GST tag for affinity purification of recombinant protein. The oligonucleotides used to construct pGEX-6P-1-Rx1(1-144) were Rx1-9 and Rx1-10.

A PCR product encompassing the full-length native *NbGlk1* cDNA was cloned into the *NcoI* and *NotI* sites of pRAP35S-YFP-4HA to make pRAP35S-*NbGlk1*-4HA. pRAP35S-YFP-4HA was initially constructed by cloning YFP into the *NcoI* and *KpnI* sites of pRAP35S. A linker encoding a *NotI* site was synthesized by annealing the oligonucleotides *NbGlk1*-7 and *NbGlk1*-8. Two copies of a 2xHA tag synthesized by annealing the oligonucleotides *NbGlk1*-9 and *NbGlk1*-10 were inserted into *NotI* and *XbaI* sites of this vector to make pRAP35S-YFP-4HA. The oligonucleotides used to construct pRAP35S-*NbGlk1*-4HA were *NbGlk1*-11 and *NbGlk1*-12. An *AscI*/*PacI* fragment from pRAP35S-*NbGlk1*-4HA encompassing the 35S promoter and *NbGlk1*-4HA fusion was cloned into the corresponding sites of the binary vector pBIN+ to make pBIN35S-*NbGlk1*-4HA. A PCR product encompassing the full-length native *NbGlk1* cDNA was introduced into Gateway donor vector pDONR207 (Invitrogen) to make pDONR207-*NbGlk1*. The *NbGlk1* gene was then recombined

into the Gateway destination binary vector pK7WGF2 (50) to make GFP-*NbGlk1*. The oligonucleotides used to make pDONR207-*NbGlk1* were *NbGlk1*-13 and *NbGlk1*-14.

GFP expression was driven from a PVX amplicon by a duplicated coat protein in pGR106 as previously described (51). An *AscI*/*PacI* fragment from pRAP35S-Rx1 (18) was cloned into the corresponding sites of the binary vector pBIN+ to make pBIN35S-Rx1. pBIN-35S-based plasmids corresponding to GFP-H2B, Rx1-GFP, Rx1-mCherry and CP106 are as described (18,29).

To make pBIN35S-Rx1-4myc and pBIN35S-Rx1-CC plasmids, the oligonucleotides Rx1-5 and Rx1-6 were annealed to make a dsDNA segment with a 5' *NotI* and a 3' *XbaI* overhang which encodes a double c-myc epitope (AAASEQKLISEEDLGEQKLISEEDLTS). This segment was introduced between the *NotI* and *XbaI* sites in pRAP-YFP to create pRAP-YFPmMyc. A 4xMyc tag was created by fusing an *AscI*-*SpeI* fragment of this plasmid with a *NheI*-*PacI* fragment from this plasmid in an *AscI*-*PacI* digested pRAP plasmid. pRAP:Rx1-4myc was created by replacing the YFP in pRAP-YFP-4Myc with the Rx1 sequence from pRAP:Rx1-GFP (18) via the *NcoI* and *NotI* restriction sites. pRAP:Rx1-4myc (Rx1 CC) was created by replacing the YFP gene in pRAP:YFPmMyc with the Rx1-CC sequence amplified with the oligonucleotides Rx1-7 and Rx1-8. Amplification with these oligonucleotides introduces an *NcoI* site overlapping the start codon and a *NotI* site immediately following the sequence coding for amino acid 144 of the CC of Rx1. The expression cassettes were transferred from pRAP to pBIN+ (52) via the restriction sites *AscI* and *PacI*.

Protein Expression and Purification - Protein corresponding to Rx1-CCNBARC was expressed from pET22b-Rx1-CCNBARC in *E. coli* Rosetta2(DE3) pLysS. A 20-mL culture was grown overnight in Luria broth supplemented with 100 $\mu\text{g mL}^{-1}$ ampicillin and 34 $\mu\text{g mL}^{-1}$ chloramphenicol at 37°C. This culture was diluted into 1 L of Luria broth supplemented with ampicillin and chloramphenicol and grown at 37°C to $A_{600\text{ nm}} = 0.7$. The growth temperature was reduced to 22°C, and protein production was induced when cells reached 22°C, for 16 h with 100 μM isopropyl- β -D-thiogalactoside. Cells were harvested by centrifugation (4,000 g, 20 minutes, 4°C). Pelleted cells were washed with 50 mM Tris-HCl, pH 8.5, 1 mM EDTA, and the pellet resuspended in 50 mM Tris-HCl, pH 8.0, 200 mM NaCl, 5 mM EDTA, 5 mM DTT, 1 % (v/v) Triton X-100. Cells were lysed

by sonication (150 s) and centrifuged (42,000 g, 60 minutes, 4°C), and inclusion bodies were washed 3 times in 30 mL of 50 mM Tris-HCl, pH 8.0, 1 M NaCl, 5 mM EDTA, 5 mM DTT, 2 % (v/v) Triton X-100, 1 M urea and twice in 50 mM Tris-HCl pH 9.0, 100 mM NaCl, 1 mM EDTA, 1 mM DTT. Sonication (60 s) was used to aid resuspension and ensure complete cell lysis. The final pellet was resuspended in 5 mL of 50 mM Tris-HCl, pH 9.0, 100 mM NaCl, 1 mM EDTA, 1 mM dithiothreitol, 8 M urea. Material was incubated at room temperature with rocking for 2 hours prior to centrifugation (42,000 g, 30 minutes, 4°C) and the pellet was discarded. The supernatant was filtered through a 0.2 µm filter, aliquoted and stored at -20°C. To refold, protein was added drop-wise slowly to a final concentration of 1 mg mL⁻¹ into 50 mM Tris-HCl, pH 8.5, 9.6 mM NaCl, 0.4 mM KCl, 2 mM MgCl₂, 2 mM CaCl₂, 0.5 M arginine, 0.4 M sucrose, 0.75 M guanidine HCl, 1 mM glutathione, 0.1 mM reduced glutathione and incubated at 4°C for 1 h. Refolded protein was centrifuged (42,000 g, 30 minutes, 4°C) and supernatant dialyzed into 20 mM Tris-HCl, pH 7.5, 50 mM NaCl, 2 mM MgCl₂ overnight at 4°C, concentrated, and stored at -20°C in 20 % (v/v) glycerol.

Protein corresponding to the Rx1 CC domain (Rx1(1-144)) was expressed from pET28a-Rx1-CC in *E. coli* BL21(DE3) pLysS. A starter culture was grown overnight at 37°C in Luria broth supplemented with 50 µg mL⁻¹ kanamycin and 34 µg mL⁻¹ chloramphenicol. The overnight culture was diluted 1:50 into fresh Luria broth with antibiotics and grown with shaking at 37°C to $A_{600\text{ nm}} = 0.8$. The growth temperature was reduced to 25°C and protein production was induced with 0.5 mM isopropyl-β-D-thiogalactoside for 3 hours. Cells were centrifuged (4,000 g, 20 minutes, 4°C). Pelleted cells were washed with 50 mM Tris-HCl pH 8.5, 1 mM EDTA and centrifuged (5,500 g, 20 min, 4°C). Cells were resuspended in twice their volume of lysis buffer (50 mM Tris-HCl pH 8.0, 200 mM NaCl, 40 mM imidazole, 5 mM β-mercaptoethanol, and SIGMAFAST™ Protease Inhibitor Cocktail Tablets). Cells were lysed by sonication (150 s) and the lysate cleared by centrifugation at (42,000 g, 60 minutes, 4°C). The supernatant was loaded onto a 5 mL HisPrep HP Ni-NTA column (GE Healthcare) on an AKTA Pure chromatography system at 2 mL min⁻¹ (GE Healthcare). The column was washed with 5 bed volumes of lysis buffer, 20 bed volumes of wash buffer (lysis buffer + 1 M NaCl), 5 bed volumes of lysis buffer, and eluted with lysis buffer supplemented with 500 mM imidazole. Peak

fractions were assessed by SDS-PAGE, pooled, concentrated, exchanged into storage buffer (50 mM Tris-HCl pH 7.5, 500 mM NaCl, 1 mM EDTA, 1 mM DTT, 20 % (v/v) glycerol), and stored at -80°C.

Protein corresponding to the Rx1 CC domain fused to GST (Rx1(GST-1-144)) was expressed from pGEX-6P-1-Rx1(1-144) in *E. coli* BL21Tuner (DE3) pRARE. A starter culture was grown overnight at 37°C in Luria broth supplemented with 100 µg mL⁻¹ ampicillin and 34 µg mL⁻¹ chloramphenicol. The overnight culture was diluted 1:50 into fresh Luria broth with antibiotics and grown with shaking at 37°C to $A_{600\text{ nm}} = 0.8$. The growth temperature was reduced to 22°C and protein production was induced with 0.1 mM isopropyl-β-D-thiogalactoside for 15 hours. Cells were centrifuged (4,000 g, 20 minutes, 4°C). Pelleted cells were washed with 50 mM Tris-HCl pH 8.5, 1 mM EDTA and centrifuged (5,500 g, 20 min, 4°C). Cells were resuspended in twice their volume of lysis buffer (50 mM Tris-HCl pH 8.0, 200 mM NaCl, 1 mM DTT, and SIGMAFAST™ Protease Inhibitor Cocktail Tablets). Cells were lysed by sonication (150 s) and the lysate cleared by centrifugation at (50,000 g, 60 minutes, 4°C). The supernatant was loaded onto a 5 mL GST-agarose column (Thermo-Fisher) on an AKTA Pure chromatography system at 1 mL min⁻¹ (GE Healthcare). The column was washed with 5 bed volumes of lysis buffer, 20 bed volumes of wash buffer (lysis buffer + 0.5 M NaCl), 5 bed volumes of lysis buffer, and eluted with lysis buffer supplemented with 10 mM reduced glutathione. Rx1(GST-1-144) was further purified by size exclusion chromatography using a Superdex 75 16/60 column (GE Healthcare) in 50 mM Tris-HCl pH 8.0, 500 mM NaCl, 1 mM DTT, 1 mM EDTA at 1 mL min⁻¹. Peak fractions were assessed by SDS-PAGE, pooled, concentrated, exchanged into storage buffer (50 mM Tris-HCl pH 7.5, 200 mM NaCl, 1 mM EDTA, 1 mM DTT, 20 % (v/v) glycerol), and stored at -80°C.

Protein corresponding to *NbG1k1*(83-402), *NbG1k1*(83-243) and *NbG1k1*(1-243) were expressed from pET14b-*NbG1k1*(83-402), pET14b-*NbG1k1*(83-243) and pET14b-*NbG1k1*(1-243) respectively in *E. coli* BL21(DE3) pLysS. Protein expression and purification was identical to Rx1-CC except that protein was eluted in lysis buffer supplemented with 250 mM imidazole.

Protein corresponding to the CAP TF was expressed and purified as described previously (53).

Yeast Two-Hybrid Analyses - Hybrigenics Services SAS (Paris, France) performed the yeast two-hybrid screen using Rx1 (amino acids 1-144) cloned into pB27 bait plasmid as a C-terminal fusion to LexA (N-LexA-Rx1-C). The screen was performed against a random-primed *N. benthamiana* mixed tissue cDNA library constructed into pP6 prey plasmid. A total of 96.6 million clones (approximately 9-fold library coverage) were screened following a mating approach with Y187 (*MATα*) and L40 Gal4 (*MATα*) yeast strains as previously described (54). To confirm protein-protein interactions, freshly transformed yeast colonies were resuspended in 1 mL sterile deionized water, and 10 µL aliquots were spotted onto medium lacking leucine and tryptophan (–L/–W) and medium lacking leucine, tryptophan, histidine (–L/–W/–H), supplemented with 10 or 50 mM 3-Amino-1,2,4-triazole (3-AT). Growth was scored after 5 to 7 d of incubation at 28°C.

Protein Binding Microarray - Protein Binding Microarray (PBM) was performed with protein derived from plasmid pTH6838-*NbGlk1*(82-244) essentially as described (55,56). *NbGlk1*(82-244) was analyzed in duplicate on two different arrays with differing probe sequences. Microarray data were processed by removing spots flagged as ‘bad’ or ‘suspect’, and employing spatial de-trending (using a 7x7 window centered on each spot) as described (56). Calculation of 8-mer Z- and E-scores was performed as previously described (57). Z-scores were derived by averaging the spot intensity for each probe containing the 8-mer, subtracting the median value for each 8-mer, and then dividing by the standard deviation to yield a distribution with a median of zero and a standard deviation of one. E-scores are a modified version of the AUROC statistic, which considers the relative ranking of probes containing a given 8-mer, and range from –0.5 to +0.5, with *E* > 0.45 taken as highly statistically significant (58). A position weight matrix (PWM) was derived using the PWMalign algorithm, which aligns the top 10 8-mer E-scores and tallies the frequency at each position to generate a PWM (56).

Gel Filtration Analysis - Gel filtration analysis of protein was performed at 4°C using an Sephacryl HiPrep 16/60 S200 HR column (GE Healthcare) on an AKTA Pure chromatography system (GE Healthcare). Protein was dialysed overnight against running buffer (20 mM Tris-HCl pH 7.4, 140 mM NaCl, 1 mM DTT). Proteins were incubated on ice for 30 min individually or together and then centrifuged at 12,000 g for 30 min. Protein loading

concentration was approximately 75 µM. Columns were run at a flow rate of 0.5 mL/min in running buffer. Thirty µL of each eluted fraction was subjected to SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and visualization with Quick Coomassie stain (Generon). Molecular weights were calibrated with gel filtration standards (BioRad): bovine thyroglobulin (670 kDa), bovine γ-globulin (158 kDa), chicken ovalbumin (44 kDa) and equine myoglobin (17 kDa).

Co-immunoprecipitation - *N. benthamiana* leaves were infiltrated with *Agrobacterium tumefaciens* strains (GV3101, MOG101) transformed with combinations of pBIN35S-*NbGlk1*-4HA, pBIN35S-Rx-CC-4myc and pBIN35S-Rx1-4myc and leaf material harvested 2 days after infiltration. 100 mg of leaf material was ground in liquid nitrogen and resuspended in 1.5 mL extraction buffer (10% (v/v) glycerol, 25 mM Tris-HCl pH 7.5, 1 mM Na₂EDTA, 150 mM NaCl, 0.6 mg/mL Pefabloc SC, 20 mg/mL polyvinylpyrrolidone, 0.1 % (v/v) Tween 20, 5 mM DTT). The supernatant was passed through a 5 mL G25 Sephadex column after pelleting the cell debris. The resulting sample was incubated at 4°C with 50 µL magnetic beads (Milenyi µMACs) and conjugated antibodies (Sigma) for 1 to 2 hours. Unbound proteins were removed by washing 5 times with washing buffer (10% (v/v) glycerol, 25 mM Tris-HCl pH 7.5, 1 mM Na₂EDTA, 150 mM NaCl, 0.15% (v/v) Nonidet P-40, 5 mM DTT). Captured proteins were released by heating beads in 1x NuPAGE LDS sample buffer (60 mM DTT). Start material (before incubation with beads), the unbound fraction and the captured proteins were separated on NuPAGE novex 12% bis-tris gels in MES buffer (50 mM MES, 50 mM Tris base, 0.1% SDS, 1 mM EDTA pH 7.3) and blotted on polyvinylidene difluoride (PVDF) membranes for immunoblot analysis. Affinity-tagged proteins were detected using peroxidase conjugated antibodies (c-Myc: goat anti-c-Myc (Abcam 9132) and donkey anti-goat peroxidase conjugated (Jackson 705-035-147), HA: rat anti-HA HRP conjugated (Roche 12013819001)). Peroxidase activity was visualized with the SuperSignal™ West Dura and Femto Substrates (Thermo Scientific) and imaged in a Syngene G:BOX Chemi HR-16 Gel documentation system.

Fluorescence Anisotropy – Double-stranded DNA substrates lacking a putative *NbGlk1*-binding site (No site), with a concatenated AGATTTC binding site (AGATTTC), and with a concatenated GGATATCC binding site (GGATATCC) were made by annealing synthetic oligonucleotides. FA-1

and FA-2 were annealed to make 'No site'. FA-3 and FA-4 were annealed to make 'AGATTTC'. FA-5 and FA-6 were annealed to make 'GGATATCC'. FA-1, FA-3 and FA-5 were end-labelled with FAM. Double-stranded DNA was annealed by mixing 10 μ M concentrations of complementary oligonucleotides in 150 mM NaCl, 15 mM NaCitrate, heating to 95°C, and cooling to room temperature over 5 hours. Changes in anisotropy were measured using a SynergyTM H4 Fluorescence Spectrophotometer (BioTek) fitted with polarizing filters (λ_{em} = 528 nm, λ_{ex} = 485 nm, bandwidth = 20 nm, averaging time = 10 s). Anisotropy was determined using 10 nM fluorescein end-labelled oligonucleotides (Eurofins MWG) with variable protein in 20 mM Tris-HCl pH 7.4, 140 mM NaCl, 1 mM DTT. Anisotropy was calculated using Gen5 software (BioTek).

Time Resolved FRET In Situ – *A. tumefaciens* strain GV3101 (pMP90) was transformed with constructs pK7WGF2 (GFP negative control), pK7WGF2-H2B (GFP-H2B positive control), pBIN35SRx1-GFP (Rx1-GFP), pBIN35S-CP106, pBIN35S-Rx1, pBIN35S-NbGlc1-4HA, pK7WGF2-GFP-NbGlc1 (GFP-NbGlc1) and experiments performed as previously (29).

Nicotiana benthamiana Hypersensitive Response Assay - *N. benthamiana* leaves were infiltrated with *A. tumefaciens* transformed with constructs pBIN35S-CP106, pBIN35S-Rx1, and pBIN35S-NbGlc1-4HA at a $A_{600\text{ nm}}$ = 0.1-0.5. Plants were grown for 4 days at 25°C with 16 h of light. Leaves were harvested, visually inspected, photographed and scored 1-5 for cell death: 1 being no visual sign of any cell death whatsoever in the infiltrated region and 5 being complete cell death throughout the infiltrated region (Figure 8C).

Overexpression Transient PVX Resistance Assay - *N. benthamiana* leaves were infiltrated with *A. tumefaciens* transformed with pGR106 with and without constructs pBIN35S-Rx1 and pBIN35S-NbGlc1-4HA at a $A_{600\text{ nm}}$ = 0.1-0.5. Leaves were grown for 4 days and then harvested. Three different 10 mm diameter leaf discs were excised and each disc was placed into a 96-well plate for each infiltrated area. The fluorescence intensity of each leaf disc was measured using a SynergyTM H4 Fluorescence Spectrophotometer (BioTek) (λ_{em} = 550 nm, λ_{ex} = 410 nm, bandwidth = 20 nm, averaging time = 10 s). An average of the fluorescence intensities for the three leaf discs was

calculated to give a value for each infiltrated area. The fluorescence intensity of each averaged area was normalised to an infiltrated area on each leaf with only pGR106.

Confocal Scanning Laser Microscopy - Imaging was performed using a Zeiss LSM 510 confocal microscope (Carl Zeiss) using settings as described (18). Agrobacteria harbouring the appropriate constructs were infiltrated on *N. benthamiana* leaves. Imaging was performed 48 hours post inoculation.

Statistical Analysis - Error bars represent either the standard deviation or standard error of the mean with the number of replicates as indicated in the legend. All replicates are independent biological experiments. Statistical comparisons (*p* values) were obtained from one-way ANOVA with the indicated post-hoc test unless otherwise indicated. *p* values in statistical comparisons are indicated in figures and indicate compared data sets as described in the figure legends. Calculated values for K_d were compared using the extra-sum-of-squares F test.

CONFLICT OF INTEREST STATEMENT

The authors declare that they have no conflicts of interest with the contents of this article.

AUTHOR CONTRIBUTIONS

M.J.C. conceived the project and wrote the paper with input from all authors. MJC, TRH, GJS, FLWT, LOP, and AG designed the experiments. CHD, PDT, EJS, OCAS, and AWHY performed the experiments.

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FOOTNOTES

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¹Department of Biosciences, ²Biophysical Sciences Institute, ³Department of Chemistry, Durham University, South Road, Durham, DH1 3LE, United Kingdom. ⁴Laboratory of Nematology, Department of Plant Sciences, Wageningen University, 6708 PB, Wageningen, The Netherlands. ⁵Donnelly Centre, University of Toronto, 160 College Street, Toronto, ON, M5S 3E1, Canada. ⁶Molecular Plant Pathology, Swammerdam Institute for Life Sciences, University of Amsterdam, Science Park 904, 1098 XH, Amsterdam, The Netherlands. ⁷ These authors contributed equally to this work

⁸ The abbreviations used are: AAA+, ATPases associated with diverse cellular activities; CC, coiled-coil; dsDNA, double-stranded DNA; FLIM, Fluorescence lifetime imaging microscopy; FRET, Fluorescence resonance energy transfer; GARP, maize GOLDEN2, ARR B-class proteins from *Arabidopsis*, and *Chlamydomonas* Psr1; GLK, Golden2-like; GST, glutathione-S-transferase; NB, nucleotide binding; NB-ARC, Nucleotide-Binding, Apaf-1, R-proteins, and CED-4; LRR, leucine-rich repeat; NLR, nucleotide-binding leucine-rich repeat; PBM, protein binding microarray; PVX, Protein Virus X; STAND, signal-transduction ATPases with numerous domains; TF, transcription factor; TIR, Toll–interleukin 1 receptor; VIGS, virus-induced gene silencing.

Table 1. Influence of hexahistidine-tagged Rx1 proteins on the dissociation constant (K_d) of NbGlk1 for DNA-binding motifs. Values are in μM ($\pm\text{S.D.}$) ND=not determined; - = no additional Rx1 protein; * $p<0.0001$ compared to No site motif; # $p>0.05$ compared to No site motif (F test).

	NbGlk1(1-243)			NbGlk1(83-243)			NbGlk1(83-402)		
	-	Rx1(1-144)	Rx1(1-489)	-	Rx1(1-144)	Rx1(1-489)	-	Rx1(1-144)	Rx1(1-489)
No site	ND	ND	ND	0.33 ± 0.02	0.85 ± 0.27	1.34 ± 0.42	0.18 ± 0.01	0.40 ± 0.02	0.42 ± 0.09
AGATTT	ND	ND	ND	$0.23\pm0.01\#$	0.76 ± 0.52	0.65 ± 0.04	$0.16\pm0.01\#$	0.46 ± 0.06	0.34 ± 0.04
GATATC	>1	>1	>1	$0.18\pm0.01\#$	0.40 ± 0.15	0.50 ± 0.03	$0.08\pm0.00^*$	0.30 ± 0.07	0.19 ± 0.04

Table 2. Influence of GST-tagged Rx1 protein on the dissociation constant (K_d) of NbGlk1(83-402) for DNA-binding motifs. Values are in μM ($\pm\text{S.D.}$); - = no additional Rx1 protein; * $p<0.0001$ compared to No site motif (F test).

	NbGlk1(83-402)	
	-	Rx1(GST-1-144)
No site	0.32 ± 0.03	1.99 ± 0.09
AGATTT	$0.22\pm0.00^*$	0.73 ± 0.07
GATATC	$0.11\pm0.00^*$	0.34 ± 0.00

FIGURE LEGENDS

Figure 1. The N-terminus of the Rx1 NLR protein interacts with the *NbGlk1* transcription factor. Yeast Two-Hybrid analysis of Rx1(1-144) bait fragment against a prey fragment of amino acids 1-371 of *NbGlk1*. Rx1(1-144) was fused to the Gal4 DNA-binding domain and *NbGlk1*(1-371) was fused to the Gal4 activation domain. Plates were grown on medium lacking leucine and tryptophan (–L/–W) and medium lacking leucine, tryptophan, histidine (–L/–W/–H), supplemented with 10 or 50 mM 3-Amino-1,2,4-triazole (3-AT). **A.** Smad vs Smurf positive control **B.** Empty pB27 bait vs empty pP7 prey negative control. **C.** empty pB27 bait vs *NbGlk1*(1-371) in prey negative control. **D.** Rx1(1-144) containing bait vs empty pP7 prey negative control. **E.** Rx1(1-144) in pB27 bait plasmid with *NbGlk1*(1-371) in pP7 prey plasmid.

Figure 2. Rx1 binds *NbGlk1* *in vitro*. **A.** Interaction of Rx1(1-144) with *NbGlk1*. On the left are representative gel-filtration chromatograms of Rx1, *NbGlk1*(83-402), and Rx1 incubated with *NbGlk1*(83-402). Peak fractions were visualized by SDS-PAGE. **B.** Interaction of Rx1(GST-1-144) with *NbGlk1*. On the left are representative gel-filtration chromatograms of Rx1(GST-1-144), *NbGlk1*(83-402), GST, Rx1(GST-1-144) incubated with *NbGlk1*(83-402) and GST incubated with *NbGlk1*(83-402). Peak fractions were visualized by SDS-PAGE.

Figure 3. Rx1 binds *NbGlk1* *in planta*. Co-immunoprecipitation of 4xmyc-tagged full-length Rx1 or Rx1-CC when co-expressed *in planta* with C-terminally HA-tagged *NbGlk1*. The labels on the Figure are as follows; Input-denotes the constructs Agrobacterium-infiltrated into *N. benthamiana* leaves; αMyc-an immunoblot probed using an anti-myc epitope tag antibody; αHA-an immunoblot probed using an anti-HA epitope tag antibody; αMyc IP-immunoprecipitation of the denoted input samples using an anti-myc epitope tag antibody; CBB-Coomassie Blue stain loading control for the denoted input samples. Immunoblot bands corresponding to Rx1-4myc, Rx1-CC-4myc and *NbGlk1*-HA from the αMyc and αHA immunoblots are indicated.

Figure 4. Influence of Rx1(1-489) on *NbGlk1* DNA-binding. Fluorescence anisotropy values are plotted against log protein concentration for DNA motifs in the presence or absence of Rx1(1-489). **A.** *NbGlk1*(1-243) with the GATATC motif. **B.** *NbGlk1*(83-402) in the absence of a specific motif. **C.** *NbGlk1*(83-402) with the AGATTT motif. **D.** *NbGlk1*(83-402) with the GATATC motif. **E.** *NbGlk1*(83-243) in the absence of a specific motif. **F.** *NbGlk1*(83-243) with the AGATTT motif. **G.** *NbGlk1*(83-243) with the GATATC motif (means ± S.E.; $n > 3$). Statistical analyses of the curves are provided in the main text and Table 1-2.

Figure 5. The influence of Rx1(1-144) and Rx1(1-489) on *NbGlk1* DNA-binding. Panels A-G shows fluorescence anisotropy values plotted against log protein concentration for varying *NbGlk1* constructs binding to different DNA motifs in the presence or absence of Rx1(1-144) ($n = 4$). **A.** *NbGlk1*(83-402) binding to DNA in the absence of a specific motif. **B.** *NbGlk1*(83-402) binding to DNA with the AGATTT motif. **C.** *NbGlk1*(83-402) binding to DNA with the GATATC motif. **D.** *NbGlk1*(83-243) binding to DNA in the absence of a specific motif. **E.** *NbGlk1*(83-243) binding to DNA with the AGATTT motif. **F.** *NbGlk1*(83-243) binding to DNA with the GATATC motif. **G.** *NbGlk1*(1-243) binding to DNA with the GATATC motif. Statistical analyses of the curves for panels A-G are provided in the main text and Table 1-2. **H.** The ratio of fluorescence anisotropy values for 10 μM *NbGlk1* or CAP binding to dsDNA with or without Rx1(1-144) (scatter plot ± S.D). The means are significantly different ($p = 0.0286$, Mann Whitney). **I.** DNA-binding for *NbGlk1*(83-402) and *NbGlk1*(83-243) was measured in the presence or absence of 10 μM nucleotide (scatter plot ± S.D; bars with different letters are significantly different ($p < 0.05$); one way ANOVA with post hoc Holm-Sidak multiple comparison).

Figure 6. The influence of Rx1(GST-1-144) on *NbGlk1* DNA-binding. The panels show fluorescence anisotropy values plotted against log protein concentration for *NbGlk1*(83-402) binding to different DNA motifs in the presence or absence of Rx1(GST-1-144) ($n = 4$). **A.** *NbGlk1*(83-402) binding to DNA in the absence of a specific motif. **B.** *NbGlk1*(83-402) binding to DNA with the AGATTT motif. **C.** *NbGlk1*(83-402) binding to DNA with the GATATC motif.

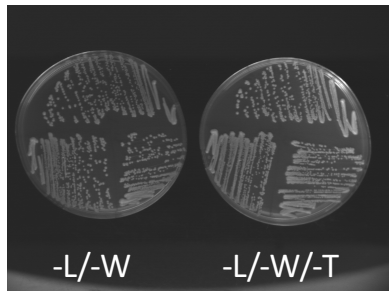
Figure 7. Binding of Rx1 and NbGlk1 protein to DNA *in situ*. **A.** Representative confocal images of nuclei and surrounding cytoplasm of *N. benthamiana* epidermal cells transiently co-expressing NbGlk1-GFP with mCherry. Images are of the NbGlk1-GFP channel (left hand panel), the mCherry channel (centre panel) and an overlay of the NbGlk1-GFP and mCherry channels (right hand panel). Co-expression was performed in the presence (upper panels) or absence (lower panels) of the p19 silencing suppressor. Scale bar represents a width of 10 μ m. Subcellular structures are indicated by N = nucleus; n = nucleolus; C = cytoplasm; V = vacuole; Ch = chloroplasts. **B.** Ratio of the long to short GFP lifetimes for a Rx1-GFP full-length construct alone and upon co-expression with NbGlk1 and the avirulent CP106 allele of the PVX CP (scatter plot \pm S.D; bars with different letters are significantly different ($p < 0.05$); one way ANOVA with post hoc Dunnett's multiple comparison). **C.** Ratio of the long to short GFP lifetimes for NbGlk1-GFP full-length construct alone and upon co-expression with Rx1 and the avirulent CP106 allele of the PVX CP (scatter plot \pm S.D; bars with different letters are significantly different ($p < 0.05$); one way ANOVA with post hoc Dunnett's multiple comparison).

Figure 8. The influence of Rx1-interactors on the immune response to PVX. **A.** The average cell death score for *N. benthamiana* leaves expressing NbGlk1 expressed in combination with CP106 and Rx1. Leaves were agroinfiltrated with *A. tumefaciens* transformed with constructs pBIN35S-CP106, pBIN35S-Rx1, and pBIN35S-NbGlk1-4HA. The pBIN35S-CP106 construct expresses only the PVX CP106 coat protein (scatter plot \pm S.D; Letters indicate statistically different data sets ($p < 0.05$) calculated using Dunnett's multiple comparisons test). **B.** A representative *N. benthamiana* leaf of the results provided in panel A. **C.** Representative images of *N. benthamiana* leaf areas for each cell death score. The number within each image indicates the assigned score for the associated degree of cell death. **D.** Ratio of fluorescence compared to a control of GFP expressed from a PVX amplicon for *N. benthamiana* leaves expressing GFP from a PVX amplicon in combination with NbGlk1 and Rx1. Leaves were infiltrated with *A. tumefaciens* transformed with pGR106 with and without constructs pBIN35S-Rx1 and pBIN35S-NbGlk1-4HA. pGR106 encodes a PVX amplicon in which GFP expression is driven from a duplicated coat protein promoter (scatter plot \pm S.D; bars with different letters are significantly different ($p < 0.05$); one way ANOVA with post hoc Dunnett's multiple comparison).

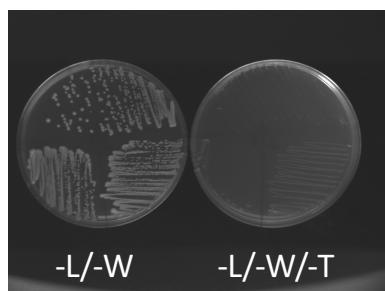
Figure 9. Model for interactions of Rx1 with NbGlk1. **A.** In the absence of PVX Rx1 interacts with NbGlk1 at DNA and a non-DNA-bound state is favoured for NbGlk1 at a non-NbGlk1 consensus DNA-binding site. **B.** Rx1-activation by PVX with a DNA-bound state favoured for NbGlk1 at a NbGlk1 consensus DNA-binding site.

Figure 1

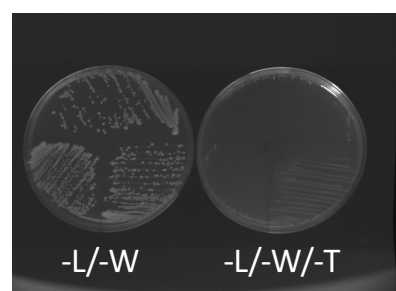
**A. pB27-Smad vs pP7-Smurf
(positive control)**



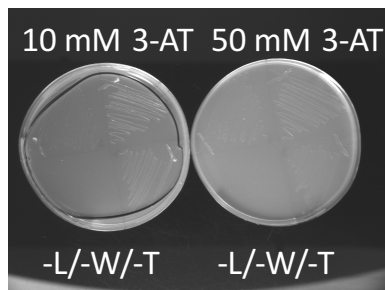
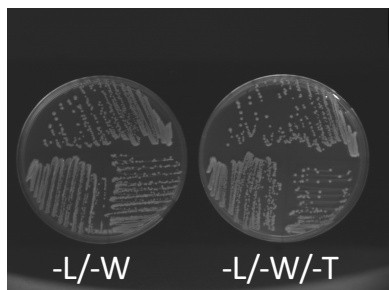
**B. pB27 vs pP7
(negative control)**



**C. pB27 vs pP7-NbGlk1(1-371)
(negative control)**



**D. pB27-Rx1(1-144) vs pP7
(negative control)**



**E. pB27-Rx1(1-144) vs
pP7-NbGlk1(1-371) (test)**

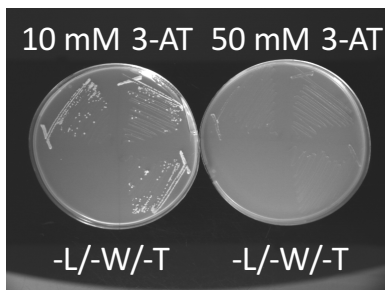
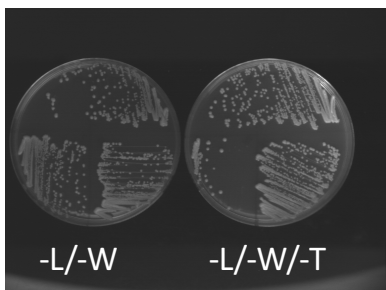


Figure 2

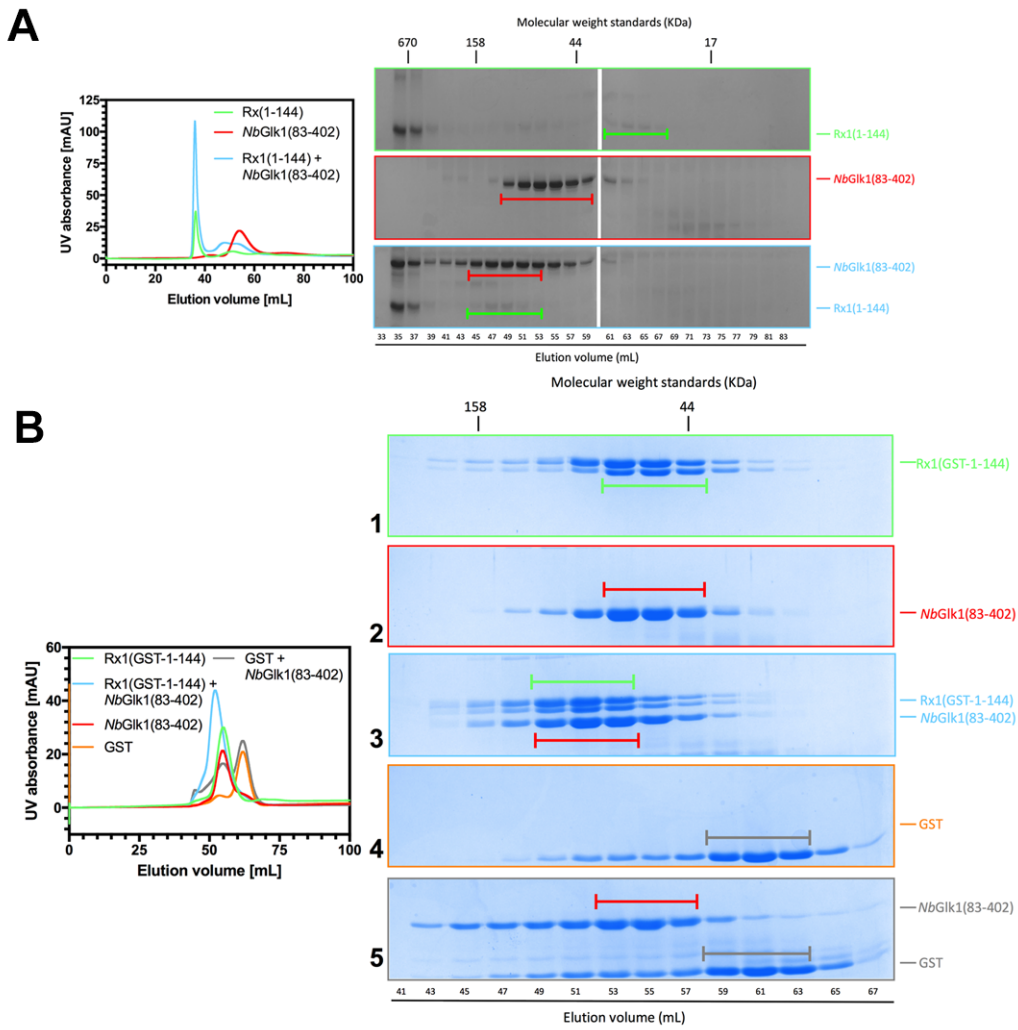


Figure 3

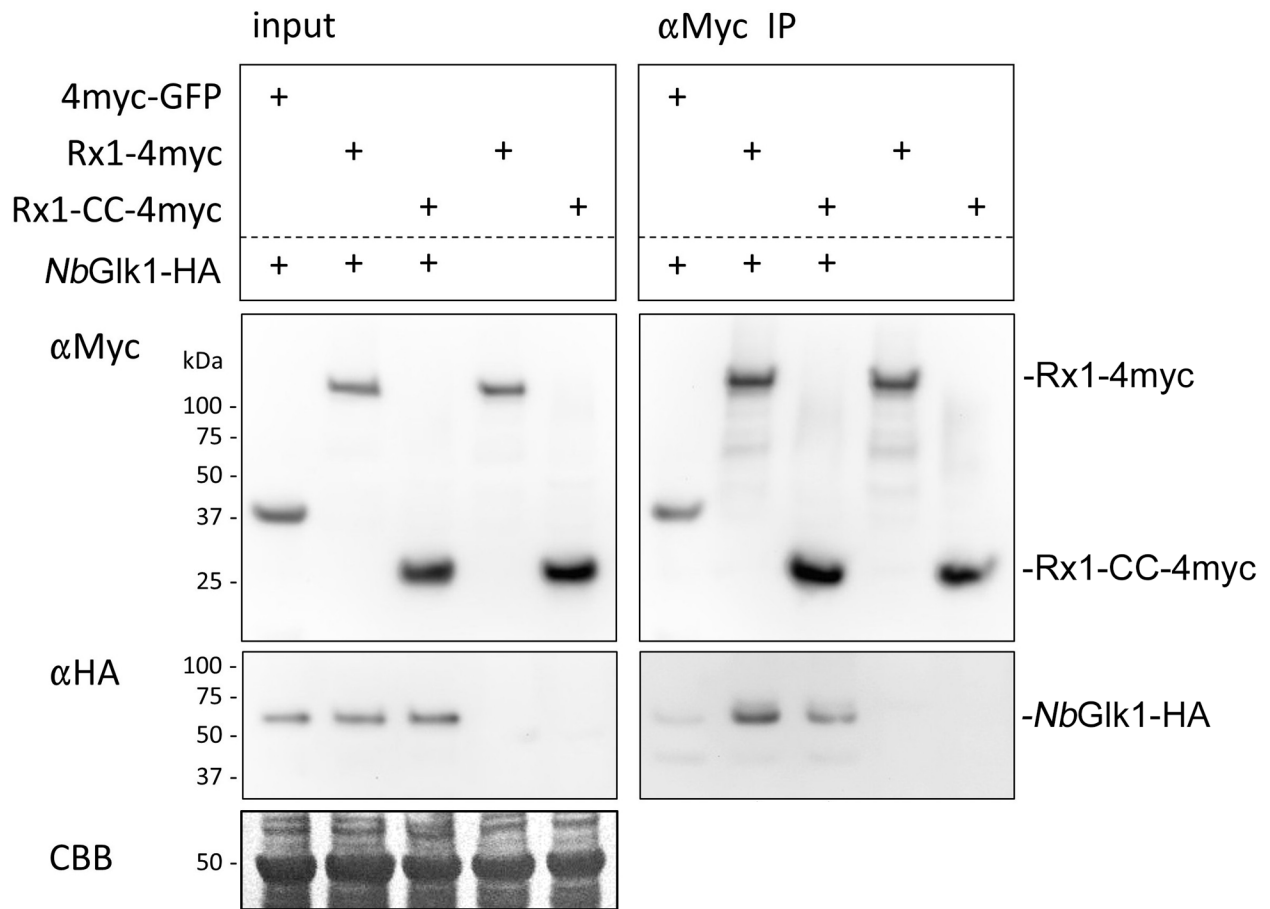


Figure 4

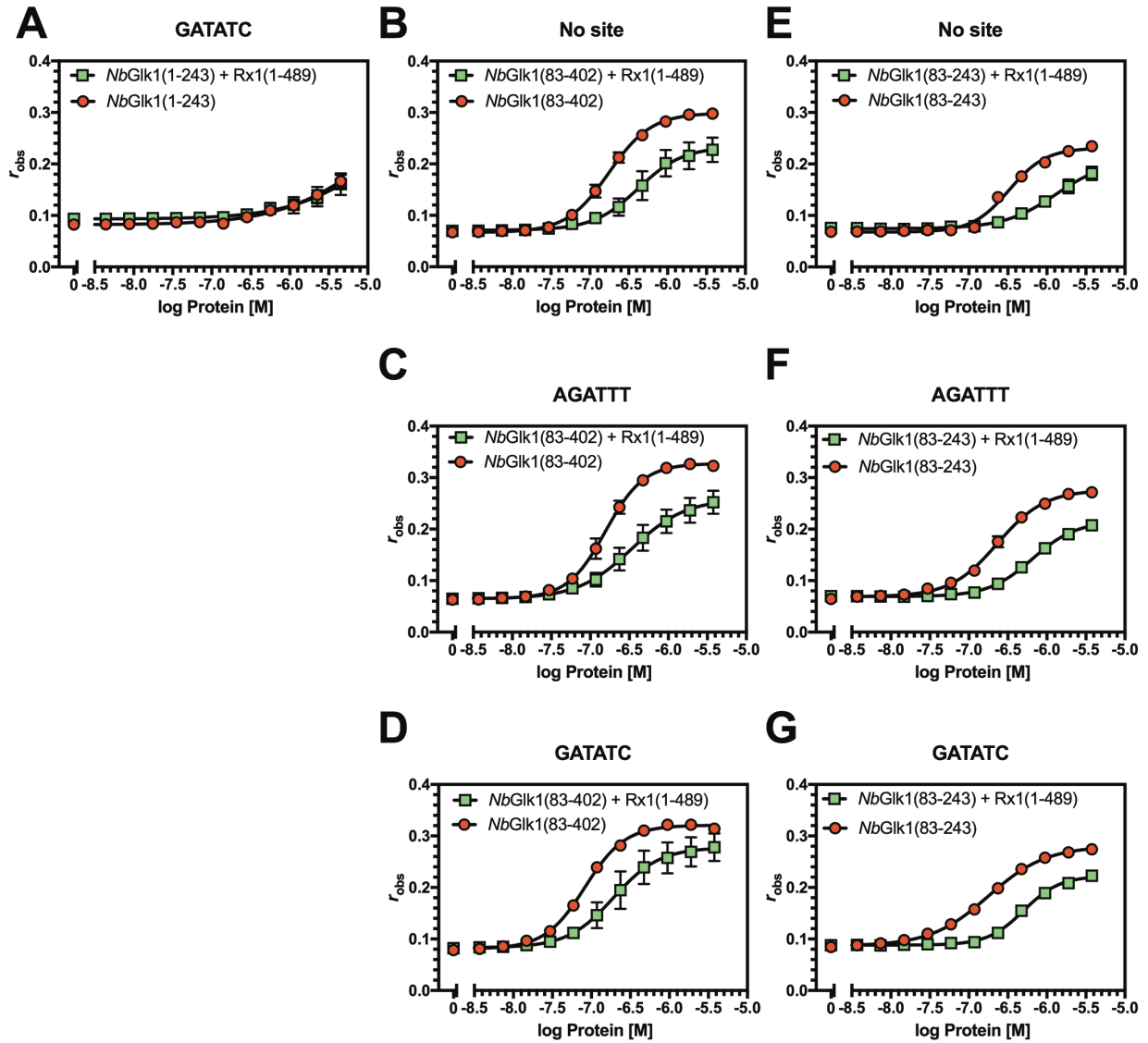


Figure 5

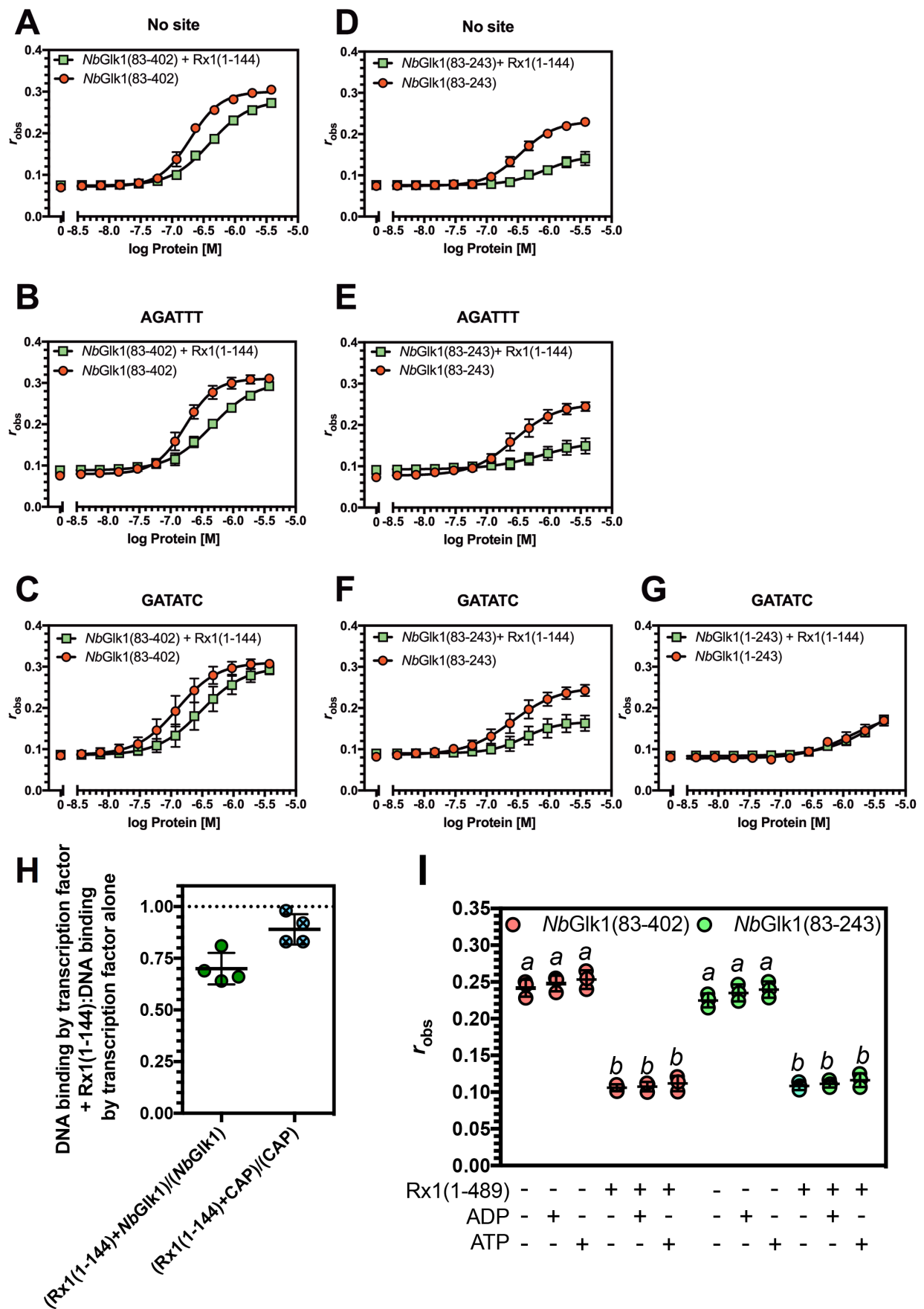
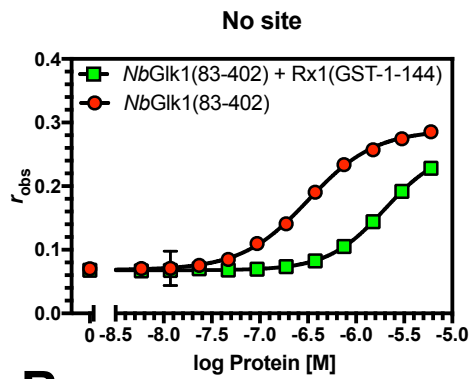
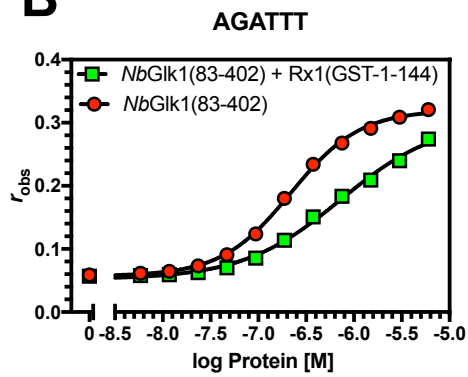


Figure 6

A



B



C

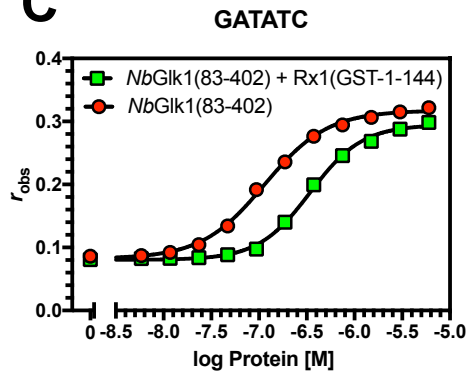


Figure 7

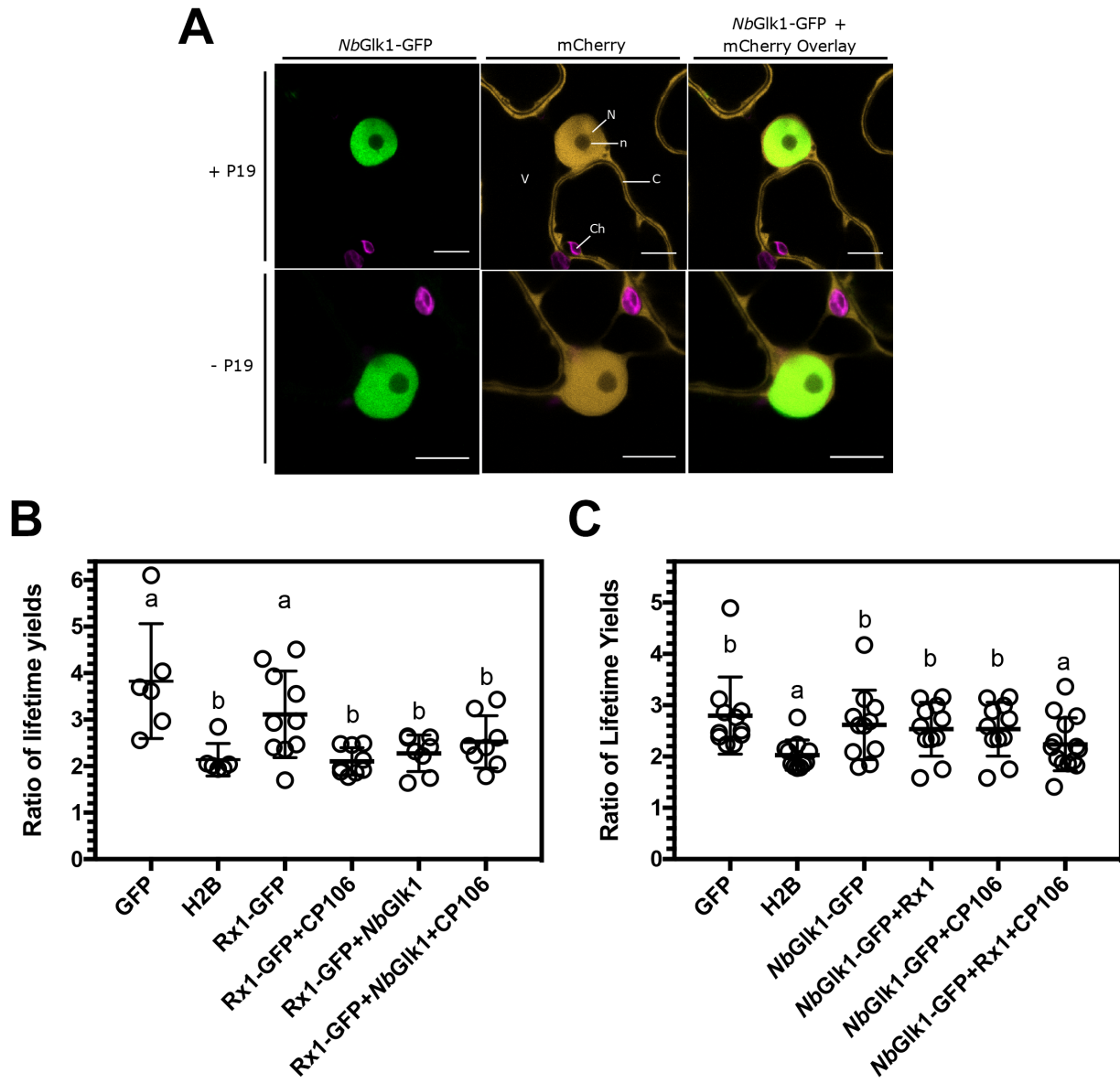


Figure 8

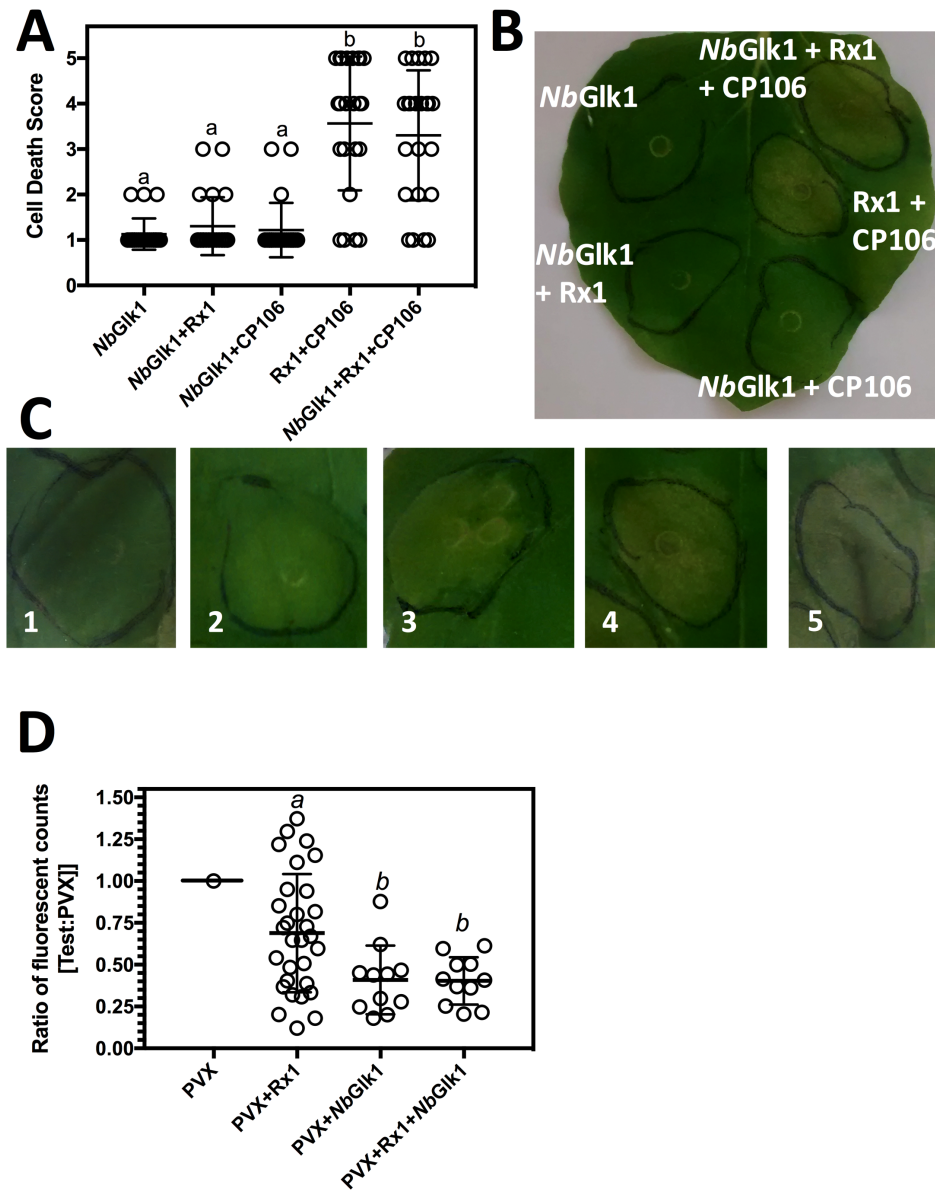
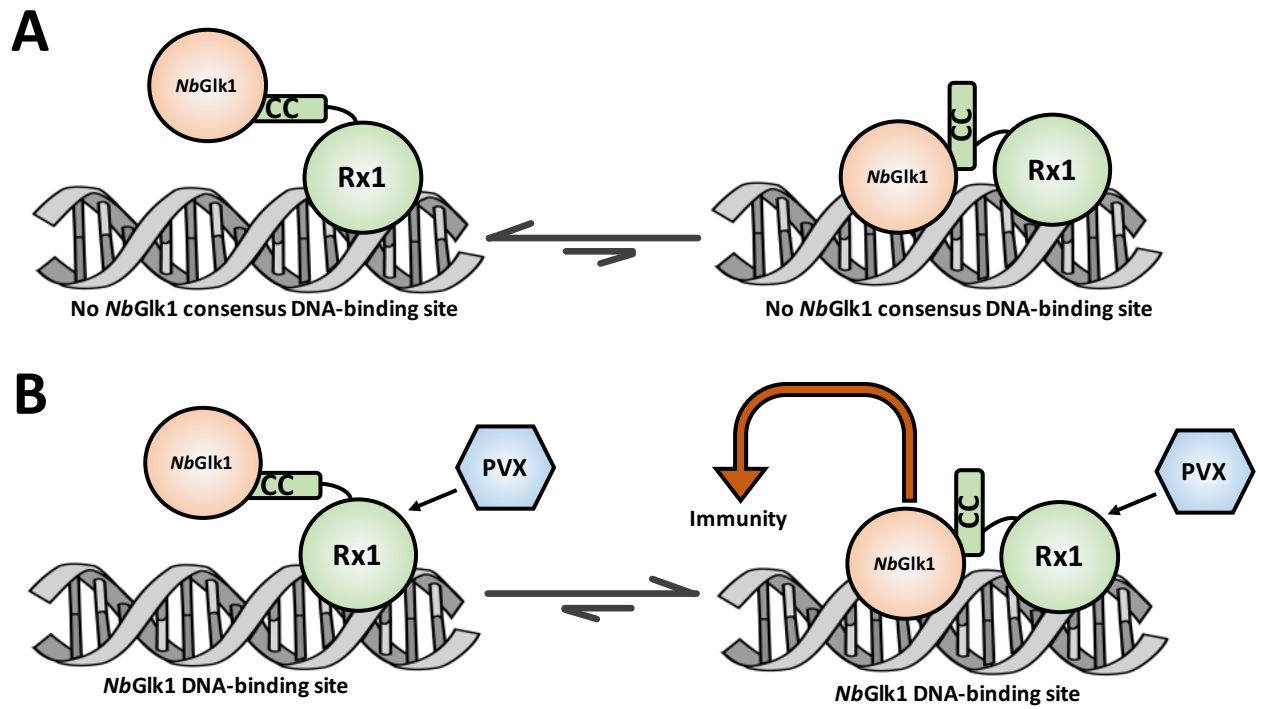


Figure 9



The intracellular immune receptor Rx1 regulates the DNA-binding activity of a Golden2-like transcription factor

Philip D Townsend, Christopher H Dixon, Erik J Sloatweg, Octavina CA Sukarta, Ally WH Yang, Timothy R Hughes, Gary J Sharples, Lars-Olof Palsson, Frank L.W. Takken, Aska Goverse and Martin J Cann

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